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Endotoxin and Immune activation in chronic heart failure: a prospective cohort study

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Summery

Background Immune ectivation in patients with chronic heart feilure may be secondary to endatoxin (lipopolysecolaride) action. We investigated the hypothesis that eltered gut permeability with beaterial translocation and endotoxeemie would be increased in patients with oederna secondary to congestive heart feilure.

Methods We compared 20 patients who had chronic heart failure with recent-onset peripheral oederns (mean age 64 years [3D 10], New York Heart Association [NYHA] class 3-3 [0-7]), 20 stable hon-oedernatous patients with chronic heart failure (mean age 63 years [19], NYHA class 2-6 [0-7]), and 14 hearthy volunteers (mean age 55 years [16]). Blochamical markers of endotocaemia, information, and immune activation were measured. Tan patients were studied within 1 week of complete resolution of oederna. Five patients survived longer than 6 months and were restudied again after remaining free of oederna for more than 3 months.

Findings Mean endotoxin conventrations were higher in cadematous patients with chronic heart failure than in stable patients with chronic heart failure (0.74 [SD 0.45] vs 0.37 EU/mL [0.23], p=0.0009) and controls (0.46 EU/mL [0.21], p=0.02). Dedensatous patients had the highest concentrations of several cytokines, After short-term diuretic treatment, endotoxin concentrations decreased from 0.84 EU/mL [0.49] to 0.45 EU/mL [0.21], p<0.05) but cytokines remelhed raised. After freshom of oxiderna for more, then 3 months after occleans resolved, endotoxin concentrations remained unchanged from the previous visit (0.49 EU/mL [0.06], p=0.45).

Interpretation Raised concentrations of endotoxin and cytokines are found in pediants with chronic heart failure during acute cedematous exacerbation. Intensified diuretic treatment can normalise endotoxin concentrations. Our preliminary findings suggest that endotoxin may trigger immune activation in petiants with chronic heart failure during cedematous oplandes.

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introduction

Some patients with chronic heart failure have features such as cardisc eacheris that may be due to activation of the immune system. Increased expression of tumour necrosis factor a (INFa) has been found in cardiac tissue of patients with chronic heart failure undergoing heart transplantation and the failing heart has been suggested as the cause of immune activation. No link between a pathogenic process and cytokine activation has been documented in human beings with heart failure or in spinal models. The cause of increased cytokine production in patients with heart failure remains unknown.

We have previously suggested that becterial endotoxin, lipopolysaccharide, contributes to immune activation in chronic heart failure. Acute venous congestion could lead to altered gut permeability for becrede, endotratio, or both, and to translocation of these menerials into the circulation, In the circulation, lipopolysaccharide is bound by a scrum protein, termed lipopolysaccharide-binding protein (LBP). The lipopolysaccharide-LHP complex can interest with the CD14 membrane protein and Toll-like algualling receptors to start a signalling cascade that leads to increased cytokine production (figure 1). The extracellular domain of the CD14 receptor is shed after interaction and serum concentrations are thought to reflect the amount of endomain and cell interaction. The lipopolysacchanide-IRP mile has been shown to be crucial for the imminosimulatory effects of lipopolysaccharide,4 High concentrations of LHP, as seen during the acute-phase response, can completely block lipopolysaccharide effects . in vitro and in a murine sepsia model,' Furthermore, patients with high concentrations of soluble CD14 (which shows endorozin-cell interaction and ahedding of CD14 from the cell membrane") have surkingly incressed concentrations of TNFa, soluble TNF receptor-1 and recepto-2, and intracellular-adhesion molecule-1.4

The degree of bowel-wall orderna cannot be directly measured. The relation between central harmodynamics and the pathophysiological features of chronic heart failure is weak. In animal models there is a poor relation between intracardiac pressures and intestinal perfusion. We therefore separated patients according to the presence or shance of a reliable marker of active venous congestion due to cardiac failure, namely peripheral orderna. Bowelwall orderna that could cause shreed gut permeability and bacterial (ie, endotoxin) translocation is most likely to occur with moderna to severe peripheral orderna.

Our main aim in this study was to measure endousin and cytokine concentrations in patients with clutteric heart failure during an acute exacerbation with peripheral orderns and after charactern and long-term treatment with direction.

Methods

Participants

We studied prospectively 14 healthy volunteers (mean age 55 years [SID 16]) and 40 patients with chronic heart follows (mean age 66 years [15], p=0-11). We did the baseline studies between April and October, 1997. 20 stable patients were recruited during outputient clinics on 3 specific days and 20 patients with medicate

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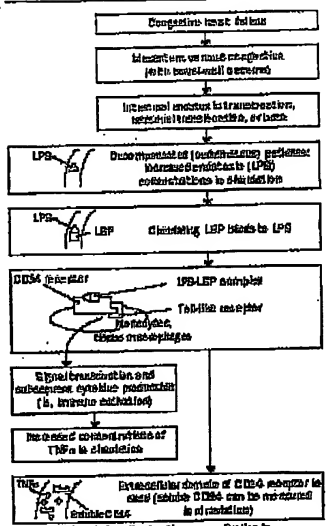


Figure 1: Endotoxin hypothesis of leanume softvation in conjustion hastifully to 173-bosohousehands.

or severe orderna represented all decompensated patients at Royal Brompton Hospital, London, UK, duting the time period, identified on attendance to the clinic or admission to the ward. Healthy volunteers were hospital stiff and relatives of parients who agreed to participate. Only one healthy person declined participation. We excluded dons from these volunteers aged participation. We excluded dons from these volunteers aged promper than 35 years to achieve a similar mean age in all groups. The causes of chronic heart failure was based on symptoms arising dislopathic dilated cardiomyopathy in 13 patients. The diagnosis of chronic heart failure was based on symptoms arising during exacults, cardiomegaly, and decommented left-ventricular dyalization (all patients had a left-ventricular ejection fraction measured by exhaustifography or radiometide ventricular particular of <00%). No patient or volunteer had clinical signs of inflation, the material attacks, or cancer.

Patients were treated with dimerics (pm38), angiorensinconverting-ensure inhibitor (p=36), dignoin (p=14), aspirin (p=17), amindament (p=16), and nimers (p=15) in various combinations. We did extended follow-up of two pedematous patients who lived close to our hospital (five New York Heart Association (NYHA) close IV, five class III) after treatment with increased doses of discretes (increase of famoranda up to 120 mg/day, with addition of bendenfumenthiselds 2-5 mg or 5-0 mg once daily, metalanne 5 mg or 10 mg once daily, or both). Of these patients, times had to be admitted for 3-8 days for intravenous discrete treatment.

After a modium of 14 days (name 7-89) we restudied these projects within 1 week of complete resolution of occluma (after treatment six NYFA class III, from thus II, mean weight loss: 3-6 kg [0-3, range 2-5-5-0]). Her patterns regarded climical

stability (one NYHA class III, four class II) and were restudied again 14-32 weeks (mean 21 weeks [7]) after the initial investigation when they had been free of peripheral occluma for more than 3 months. The remaining five patients did not entain a stable clinical state and died 2-8 months after the initial investigation without having been remained. The research protocol was approved by the other committee of the Royal Brompton Hospital, and all patients and volunteers give written informed constant.

Assavs

Blued samples were collected after rest for at least 15 min. A polyshylene eatheter was interned into an anteenhind wein and 8 mL of blond were drawn into endotoxin-free tubes (Endo Tube RT, Chromogenix AB, Sweden). 30 mL samples were also taken for blochemical and cytokine measurements. After immediate contribugation, blood and plasma samples were stored at —80°C until analysis. In addition, 5 mL blood was taken into tubes containing effects acid for fivorescence-activated cell-sorting analysis.

Concentrations of endotosin were measured with a commercially available hit (Lincolns Amehotyte Lyunie QCL-1000 test kit, Blowhitaker line, Walkersville, USA). The normal concentration of codetosin in this array in healthy people is <0.50 BU/mL. The within-easy coefficients of variation at concentrations of 0.35 BU/mL and 0.82 BU/mL were 9.9% and 9.6% between easy coefficients of variation were 16.8% and 13.3%, respectively. For repeated blood samples in non-ordernatous patients the coefficient of variation was 10.8%. The lower limit of describin was 0.03 BU/mL.

LEP was measured by HISA. Total TNFa was measured with an ELISA kir (Medgenix, Flancus, Belgiana sensitivity 3-0 pg/mL; text not influenced by soluble TNF receptors). HISA kip (R&D Systems, Minnespolis, MN, USA) were used measure soluble TNF receptor-1 and receptor-2, and invested in 6; hower limits of detection of the assays were 25 pg/mL, 2 pg/mL, and 0-0094 pg/mL, respectively. Soluble CD14 was satessed by HISA (HIL, Hamburg, Germany). Planca proceduration concentrations were measured by on immunical managements.

(BRAHMS, Berlin, Germany).*

In a subgroup of ten non-nedemators and seven nedemators patients, as well as in all healthy volunteers, whole-blood samples were taken in putsusion edem-acid tubes (Vucutainer Systems, Paleon HD, Oxford, UK) and stained with finorescently labelled monuclous! smillodies (Coalier Electronics, Lamon, UK) to

	Healthy with mirrors (mails)	(Hasti) Calenta Calenta	CRIF ondersa (11420)
Dersoglandsy (meson (SD)) Age (poten) Wasels (ke)	56 pus) 74 (7)	53 (19) 76 (9)	78 (E) 64 (E,O)
WYNA class		26(0-7)	8-3 (D-E)*
Cracco (schembs Ellopethic clients	•••	15 4	11 9
Control (mason (SDP) Social (masol/U) Crestition (masol/U) Una (masol/U) Una cost (masol/U) Aspendes underterminate (U/U)	230 (145) 82 (15) 54 (104) 804 (181) 28 (141)	557 (4-5) 531 (55) 11-0 (7-6) 417 (245) 24 (5)	184 (4-87) 229 (187) (5 20-7) (12-4) (5 640 (17-4)*) 23 (8)
North animaliania (V/L)	28 (LD)	TEN EL	24 (4 <u>)</u> []
tymphocyte profile (ypman (SD)) CD4 CD3 CD4/8 mile CD4/25 acto GD6/25 sate	47 (E) 22 (7) 2-3 (1-2) 5-7 (4-2) 4-7 (2-4)	51 (T1) 28 (14) 83 (24) 5-5 (2-4) 8-7 (4-6)	35 (15)) 28 (20) 20 (18-7) 21-6 (10-7)] 21-6 (10-7)]

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*pro-OCI vs Cit- no nedama. †pro-OCI vs heality substituena. †pro-OCI vs heality
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Teble 1: Churacteristics of patients with chronic heart failure with and without peripheral opening and healthy volunteers

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	Haveley volunts are	Gif no outsink	Crif contains
	(n=14)	(=20)	(n=20)
Endotesin (EL/mL) Life (rg/mL) Lipopolys sociacióe/log	0-45 (0-21)	0-37 (D-23)	0-74 (D-45)*†
	8-8 (4-5)	10-4 (D-3)	12-1 (B-0)
	0-64 (0-20)	D-44 (D-30)	0-75 (O-48)‡
THE (PE/ML) Solutio THE receptors.	24-8 (9-6)	96-8 (1/9)	95-5 (12/3)†\$
	708 (21%)	1077 (5 29)	9822 (1390)†\$
(pg/ml.) Salable TNF repaparor (eg/ml.)	1465 (885)	2018 (1280)	81A3 (1690)35
Solgia CDIA (m2mi.)	2455 (563)	35)4 (454)	4249 (588)14
Promiciporio (m2/mi.)	87 (15)	108 (73)	1A5 (34)
(ntertaulino (pg/mi.)	2-0 (0-4)	4-3 (5-5)	14-7 (17-3)19
Granctico protein (m2/l.)	5-6 (1-7)	8-9 (3-5)	13-7 (17-137)

*pcOdly to healthy voluntains, [pcOdl) as CHF no anderna, \$pcOdl as CHF no orderna, \$pcO-OS as lightly voluntains. \$pcO-OS we healthy voluntains. \$pcO-OS we character as \$pcO-OS we call the performs.

Table 2: Neon (50) plasma concentrations of endotude and inflammatory markets to healthy volunteers and patients with chronic heart fellows

determine peripheral lymphotype phenotype and the proportion of CD25 receptor-positive T cells. A saining excess of surbody, determined by furnism (data not shown), was placed into 12×75 mm polymorphene tubes (Ethay, Hampahim, UK). Two tubes were sunlysed for each parient's sample. The first tube contained council mouse antihuman santibodies isotopically matched to the test subodies in the stood tube. The sunbody-functionate employates tastd were CD3-PC3, CD4-FITC, CD8-BCD, and CD25R-RD1. The functi-acid lysed whole-blood protocol was used in the multi-Q-prep (Coulter Ricetomics, Laton, UK). Lymphotype gating was set on forward compated with side-scatter dut plot, and competantism was established by the combining of single-enjour-strined lemonyte populations. Four-enjour flow symmetry was done on the Conker XI-MCL with Symem II anforms (version 2.0).

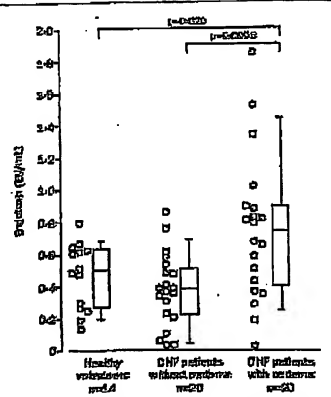
Statistical analysis

We satested normality of distribution with the Holmogorov-Smirnow test. Copaired Student's a nest, petrad t test, ANOVA with Faher's port-hoc test (with allowance for multiple testing), and the Mann-Whinty U test were used where appropriate. Data are presented so mean (SD). We size used universign-correlation and prelivariate-correlation analysis to establish the relation between variables. We mak p<0.05 to be algorithm.

Results

In tables 1 and 2, baseline clinical characteristics and results of immunological and humoral measurements are shown. Endotonin communications were highest in heart-failure patients with peripheral orderna, compared with heart-failure patients without orderna (98%) and controls (59%, p=0-0027; figure 2). Planus concentrations of LHP did not differ between groups, but there was a raised lipopolysaccharidelog LHP ratio in the heart-failure patients with orderna compared with those without orderna (71%, p<0-01). In ordernations heart-failure patients, plasma communicates were algorithmistly higher for C-reactive protein, TNFa, subble TNF receptor-1 and receptor-2, interdenkin-6, and soluble CD14 (vable 2) than for all other groups.

Among all perticipents (n=54), concemnations of subble CD14 courdened significantly with endotonin (r=0-30, p<0-05). This correlation was not significant when patients or healthy volumeons were analysed separately. In all patients with chaotic heart failure, boluble CD14 correlated with TNFs (r=0-32, p<0-05) and soluble TNF receptur-1 (r=0-45, p<0-01). There was a correlation between soluble CD14 and soluble TNF neceptur-2 in patients with stable chaotic heart failure (r=0-61, p<0-01).



Agure 2: Pleans emission concentration in healthy volunteers and health flure patients with and without cedema. Short herizontal lines—10th and 90th percentiles; long herizontal lines—20th, and 75th percentiles.

No simple correlations existed between creatinine or uses plasma concentrations and lipopolyseccharides at buscline, nor between changes in maximus of kidney function over time compared with changes of lipopolyseccharide or cytokine concentrations over time (data not shown). Therefore, a bias because of latent shootmalities of kidney function seem in some ordernators patients is unlikely.

Intensive discrete treatment for a mean of 23 days (8) in ten patients with change heart failure resulted in a mean weight decrease of 3.6 kg (range 2.5-5.0), and improvement in the functional NYHA class in time of the ten patients. In eight of these, endotatin plasma concentration was decreased from 0.96 HU/mL (0.47) to 0.45 HU/mL (0.24). In two patients with normal concentrations of endotation after discrete treatment were 9% and 36% higher than at baseline, but still in the normal range (<0.5 HU/mL). In all ten patients the lipopolysechanide concentrations fell from 0.84 HU/mL (0.49) to 0.45 EU/mL (0.21, p=0.049;

	Busins (p=10)	After d'apolis transment (polis)	Р
Endotreth (ELI/ml.)	0-84 (0-49)	0-45 (0-21)	40-03
LEP (DEPOL)	10-3 (2-7)	12-7 (7-6)	0-27
Upopolysacehtridt/log LEP ratio	0.86 (0.44)	D-28 (D-67)	0-039
Tigge (pg/tgl.)	394 (\$\$/Z)	40-2 (12-0)	0-82
Solnible THF receptor4. (pg/mL)	2520 (1314)	का स्ट्रा १७३८मी	6-09
Soluble TNF receptor-2 (or/mi.)	8751 (1185)	4029 (1ABT)	0-46
Exploide CO14 (ng/cd.)	4474 (592)	4430 (764)	0.69
Proceedings and and and a	153 (EZ)	210 (104)	0-17
Hartedon-G (re/ml)	194 (294)	18-3 (23-9)	D-60
Consider protein (mg/L)	196 (120)	20-0 (20-7)	0-00

Table 8: Mass (SD) plasma concentrations of endotroin and inflammatory markets in occumatous policula before and affect duratio treatment

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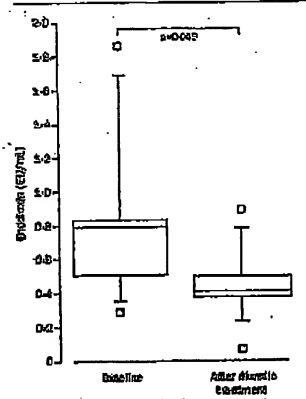


Figure 3: Effect of Interested disselfs treatment on phisima emilologic concentrations in ten patients with strends heart failure who had prophesal codema. Short hetcortof free-tith and 20th persentiage larg borbonial librar-25th, 50th, and 76th persentiage orderes votes extend 20th orderes who extending outside 3.0th and 80th persentities.

figure 3). The effect of directic treatment on the endominal and inflammatory markets are shown in table 3. During extended follow-up, five patients were restudied when free of oederns for more than 3 months after 21 weeks (7). Endouvin concentrations at the third visit did not differ from those at the second visit after a mean of 19 days (0-39 [0-22] to 0-49 HU/mL [0-06], p=0-45), but TNFs concentrations were lower (39-6 [12-4] at 31-0 py/mL [5-7], p=0-079)

Discussion

We have shown that endotoxin concentrations and proinflammatory cytokines are raised in patients with heart failure who have peripheral orderna. Refeed endotorin concentrations were normalised by lengthened divertic treatment. The endommernia in these patients was not associated with a strong acute-phase response that would have led to an increased hepatic LBP synthesis and subsequent blocking of lipopolysactharide effects. These results lend credence to the hypothesis that bacterial endotoxin may be an important stimulus of immune activation in patients with chronic heart finance. This finding may open various options for treatment of patients with chronic hear failure that could be directed against becteris in the bowel, the musicestion process, and endomain itself, the binding sizes of becaraist endomain on immone competent cells, or both.

The complex of endotoxin and endotoxin-binding protein serious monocytes and that a macrophages via the CD14 and Toll-like receptor proteins. Which stimulates the production of TNFs and other cytokines (figure 1). Previous studies suggested that increased soluble CD14 concentrations might be related to endotoxacmia. We

established that ordenatous parieties have the highest concentrations of soluble CD14 and lipopolysaccharido, but in homogeneous groups of patients there was no direct numeric relation between the two variables. Shed, and therefore soluble, CD14 receptors are thought to reflect the amount of endotoxin/cell interaction in the long term. By contrast, endotoxin has a short plasma half-life (10-50 min), which may explain why soluble CD14 concentrations are more closely related to cytoking than endotoxin concentrations.

The concentrations of endotonin in our study were well below those seen in septic shock." Patients with change heart failure had no signs of active infection, and the moderate increases in plasma endotoxin are in keeping with the hypothesis of a translocation process. Possibly, it is endotoxin itself rather than bacteria that translocates. Lipopolysacthatides at baseline did not correlate significantly with renal function (as estimated by creatinine and uses) although this finding cannot completely exclude an influence of renal function on cytokine elegance.

Although immediated diagetic therapy resulted in normalisation of endotoxin concentrations, tresument did not lead immediately to lowered cytokine plasma concentrations, which is in beeping with a previous study.15 This effect may be due to a concentration effect, resulting from the loss of up to 5 kg body water or long-term activation of monocytes or macrophages after brief exposure to an endotoxin minulus during a phase of clinical deterioration with increased venous congention. Alternatively, the lack of cytokine decrease immediately after clinical improvement may be due to a change in monocyte or mecrophage lipopolysaccharide sampiwity (ie, nomination endomin concentrations may still cause incremed symbine production), Indeed, such an increased cellular sensitivity to lipopolysaccharides has been documented in patients with chronic heart follow who had scare decompensation.19 The previously documented raised TNFa concentrations in cardiac tissue of patients with endstage chronic heart failure may also be due to cardiomyscytes or tistue monocytes releasing increased of cytokines upon stimulation ipopolysuccharides because of decompensation Œ hypersenshive cardiomyocytes. In cardiomyocytes of heart transplantation recipients (especially in patients with ischeemic chronic heart failure) increased baseline and Hopolyspecharide-stimulated TNF a production has been reported." In our study, after a long phase of clinical stability, TNPs planus concernations showed a strong trend to decrease back to normal, and, therefore, the process of normalization of cymbine secretion actins to be slow.

Tolerance of monogres or macrophages to endotoxin can be induced in vivo and in vitro by endotoxin itself. Such an effect frequently occurs after severe injury." One important mediator of lipopolysacthanides hyposenshivity is interieshin-10." Compared with countrils, we found interleukin-10 to be lower in stable patients with chronic heart failure. Increased cardiac-wall stress and general time hypoxia (both via local free-ordical generation and subsequent stimulation of the mulear factor-x B pathway) and hormonal catabolic and anabolic imbalance (especially in patients with muscle wasting.") may lead to immunological hypersensitivity. Endotoxin may, therefore, be an important stimulation of the problem production in the heart and in the periphery even in the absence of ordenta. In-vitro low concentrations of lipopolysaccharides have

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deminantal effects on cardiomyocytes. These effects are indirect through the release of other substances, a but direct effects also have been seen." In vivo there may be a dynamic belance between heart function and immuno emplation in parients with chronic heart failure, and over time patients with frequent ordenatous episodes may deteriorane because of the cardiodepressant and metabolic concernences of raised TNFn concentrations. Better control of ociena in chronic heart follow may therefore, be beneficial.

In stable ambulatory periants with chronic heart failure, a significant excess concentration of cytologica from the heart could not be shown," which suggests that cardiac production may not be the main source of the raised peripheral cytokine plasma concentrations. In support of the importance of peripheral hypoxia, measures of increased oxidative stress have been found to correlate with soluble TNF receptor-1 and receptor-2 concentrations," We have shown that peak leg blood flow after ischaemia in clinically stable patients with chronic heart follow is inversely related to TNFa plasma concentrations. This effect may be due to a relation between hypoxia and TNFa production or toxic effects of TNFa on endothelial function." Hypoxia may not be the most important cytokine trigget in chronic heart failure because of the planna profile. Raised ionerleukin-6 emembrations can be stributed to peripheral hypoxic conditions," which will occur in chronic heart failure, but there is no report that hypoxia leads to mised concentrations of TNFc, procelemnin, or soluble TNF receptor-1 or receptors are, by comment, characteristic of endomain action, but not of hypoxic disorders.2

This study shows the presence of salect plasms endomein concentrations in patients with chronic heart fallure and peripheral oederns. In the presence of unchanged concentrations of endotosin binding protein, the raised endotusin contentration reflects a potentially pathogenic situation that leads to cytokine induction. We show that normalisation of endotoxin concentrations can be achieved by intensified distrain treatment. Bacterial endoudn may be an important athenths of immunit activation in patients with chronic heart failure. Our studies are preliminary and further investigations are needed, Nevertheless, these findings may open various new options for treatment directed against bacteria in the bowel, the manelocation process, and endottenin itself, the binding sites of bacterial endorsain on income competent cells, or both.

Contributors

Compositions
Surfan Andrew Construction of the auditorial hypothesis and
with Philip Profit—Wilson designed the study, Josef Nilchaust and
Surfan Arber coordinated the study. Josef Nilchaust and
successes with the help of Mathies Reachings. Michael Keung measured
cudetonia and TNFs. Match Duminguez did all lymphocyte studyes.
Hans-Direct Volk and Raif Behannun selvined on instancionical issues and resoured all other symblers and LEP. Josef Nichauer and
Suffer, American des des and preparted the manuscript with the help
of Philip Pools-Wilson and Andrew Coses. Hem-Dieser Volk and
Ref Schemens, edited the manuscript.

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